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			ART UNIT	PAPER NUMBER
		DECENT	1632	13
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Please find below and/or attached an Office communication concerning this application or proceeding.

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BY:

OIP TO				
MAR 0 1 2003		Application No.	Applicant(s)	
Office Action Sum		09/900,700	ALLEN, KEITH D.	
Gruce Acron Sum	mary	Examiner	Art Unit	
		Peter Paras, Jr.	1632	
· · · · · · · · · · · · · · · · · · ·			ith the correspondence addr	ess –
A SHORTENED STATUTORY F THE MAILING DATE OF THIS C - Extensions of time may be available under after SIX (6) MONTHS from the mailing dat - If the period for reply specified above is less - If NO period for reply is specified above, the - Failure to reply within the set or extended p - Any reply received by the Office later than the earmed patent term adjustment. See 37 CFI Status	the provisions of 37 CFR 1.136 e of this communication. I that thirty (30) days, a reply veraximum statutory period will priod for reply will, by statute, cores months offer the section.	(a). In no event, however, may a within the statutory minimum of thin apply and will expire SIX (6) MO	reply be timely filed ty (30) days will be considered timely. VTHS from the mailing date of this comm	nunication.
1) Responsive to communication	ation(s) filed on <u>09 Oc</u>	tober 2002 .		
2a) ☐ This action is <b>FINAL</b> .		action is non-final.		
3) Since this application is in closed in accordance with Disposition of Claims	condition for allowan	ce except for formal ma		
4)⊠ Claim(s) <u>1-23</u> is/are pendii	ng in the application.		RECEIVED	)
4a) Of the above claim(s) <u>1</u>		re withdrawn from cons	ideration MAR 0 7 2003	
5) Claim(s) is/are allow	ed.			
6)⊠ Claim(s) <u>8,10 and 17-22</u> is/	are rejected.		TECH CENTER 1600/2	<b>30</b> 0
7) Claim(s) is/are object				
8) Claim(s) are subject	to restriction and/or e	lection requirement.		
Application Papers				
9) The specification is objected				
10)⊠ The drawing(s) filed on <u>06 Ju</u>	<i>ıly 2001</i> is/are: a)⊠ a	ccepted or b) objected	to by the Examiner.	
Applicant may not request that	at any objection to the d	awing(s) be held in abeva	nce. See 37 CER 1.85(a)	
11) I he proposed drawing correct	ction filed on is	: a)□ approved b)□ di	sapproved by the Examiner.	
If approved, corrected drawin	gs are required in reply	to this Office action.		
12) The oath or declaration is ob		iner.		
Priority under 35 U.S.C. §§ 119 and				
13) Acknowledgment is made of	f a claim for foreign pr	iority under 35 U.S.C. §	119(a)-(d) or (f).	
a) ☐ All b) ☐ Some * c) ☐ No	one of:	_		
1. Certified copies of the	priority documents ha	ave been received.		
2. Certified copies of the	priority documents ha	ave been received in Ap	plication No.	
3. ☐ Copies of the certified	copies of the priority	documents have been r	eceived in this National Stag	је
14) Acknowledgment is made of a	claim for domestic or	iority under 35 U.S.C. &	110(a) (ha	
a) ine translation of the for	eign language provisi	onal application has been	n received	lication).
15) Acknowledgment is made of a	claim for domestic p	iority under 35 U.S.C. 8	& 120 and/or 121	
Attachment(s)	•	, 3	3 GITGEOF 121,	
1) Notice of References Cited (PTO-892)		4) Interview Su	mmary (PTO-413) Paper No(s)	
Notice of Draftsperson's Patent Drawing F     Information Disclosure Statement(s) (PTO	Review (PTO-948)	5) L Notice of Inf	ormal Patent Application (PTO-152	<u> </u>
U.S. Patent and Trademark Office	- 1779) raper No(s) <u>3</u> .	6)		_
PTO-326 (Rev. 04-01)	Office Action	Summary	Part of Panor	

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#### **DETAILED ACTION**

#### Claims 1-23 are pending.

## Information Disclosure Statement

The IDS filed on 10/29/01 has been considered. It appears from the file wrapper of the instant application that there may have been an additional IDS submitted on or around 11/28/01 as Paper No: 4. However, the additional IDS has not been found by the Examiner and cannot be considered at this time. If the additional IDS becomes available or is resubmitted by Applicants then it will be considered by the Examiner.

#### Specification

The Brief Description of the Drawings in the instant specification is objected to because there is no description of Figure 2A.

## **Sequence Compliance**

This application contains sequence disclosures that are encompassed by the definitions for nucleotide and/or amino acid sequences set forth in 37CFR 1.821(a)(1) and (a)(2). However, this application fails to comply with the requirements of 37 CFR 1.821 through 1.825 for the reason(s) set forth on the attached Notice To Comply With Requirements For Patent Applications Containing Nucleotide Sequence And/Or Amino Acid Sequence Disclosures. Figure 2A comprises an unidentified sequence.

Applicants are required to comply with all of the requirements of 37 C.F.R. §§

1.821 through 1.825. *Any* response to this Office Action, which fails to meet all of these

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requirements, will be considered non-responsive. The nature of the noncompliance with the requirements of 37 C.F.R. §§ 1.821 through 1.825 did not preclude the examination of the application on the merits, the results of which are communicated below.

#### Election/Restrictions

Applicant's election with traverse of Group III, claims 8,10, and 17-22, in Paper No. 12 is acknowledged. The traversal is on the ground(s) that the Examiner has not shown that a serious burden would be required to examine all the claims. This is not found persuasive because each of the Inventions requires a separate search status. In particular, it is maintained that the products of Groups I, II, III, VI and VII are different each from the other; they each have different chemical structures and can be used in materially different methods that require different technical considerations. For example, the DNA targeting construct of Group I can be used to disrupt a CRFR2 gene in a somatic cell in vitro, The cells of Group II can be used to produce and isolate a protein in vitro, the transgenic non-human animal of Group III can be used as a model of disease, the unknown agent of Group VI can be used for modulating the expression of CRFR2 in a somatic cell in vitro, and the phenotypic data of Group VII can be used for statistical analysis with a computer. It is maintained that the products of Inventions I, II, III, VI and VII are distinct due to their divergent subject matter (DNA targeting construct. cell s, transgenic non-human animal, unknown agent that can modulate the expression of a CRFR2, and data in an electronic database) and are separately classified and searched.

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It is maintained that the methods of Groups IV and V are distinct, comprising different methodologies and using different products. For example, the method of Group V can be practiced in a somatic cell *in vitro*, while the method of Group IV is required to be practiced in a transgenic non-human animal. It is maintained that the methods of Groups IV and V are distinct as they are directed to different methods that require the use of different products that need different technical considerations (transgenic non-human animals and somatic cells *in vitro*) and are separately searched and classified.

It is maintained that the products of Groups I, II, III, VI and VII are distinct from the methods of Groups IV and V; the products of Groups I, II, III, VI and VII can be used in methods, which require different reagents and technical considerations from the methods of Groups IV and V. For example, the DNA targeting construct of Group I may be used as a probe in a hybridization assay *in vitro* while the transgenic non-human animal of Group III may be used to produce antibodies to an antigen, the cells of Group II can be used to produce a protein *in vitro*, while the method of Group V may be used to identify agents that modulate the expression of a CRFR2. The method of Group IV may be practiced with agents that have different chemical structures from the agent of Group VI. It is maintained that the products of Groups I, II, III, VI, and VII are distinct from and can be used in different methods (hybridization assays, generating antibodies, producing a protein) from the screening methods of Groups IV and V.

Therefore it is maintained that all the inventions are distinct each from the other for the reasons given above. The requirement is still deemed proper and is therefore made FINAL.

Please note that after a final requirement for restriction, the Applicants, in addition to making any response due on the remainder of the action, may petition the Commissioner to review the requirement. Petition may be deferred until after final action on or allowance of claims to the invention elected, but must be filed not later than appeal. A petition will not be considered if reconsideration of the requirement was not requested. (See § 1.181.).

Claims 1-7, 9, 11-16, and 23 are withdrawn from further consideration pursuant to 37 CFR 1.142(b), as being drawn to a nonelected Invention, there being no allowable generic or linking claim. Applicant timely traversed the restriction (election) requirement in Paper No. 8.

## Claim Rejections - 35 USC § 112, 1st paragraph

The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

Claims 8, 10, and 17-22 are rejected under 35 U.S.C. 112, first paragraph, because the specification, while being enabling for a transgenic mouse whose genome comprises a homozygous disruption of the nucleotide sequence set forth in SEQ ID NO:

scope with these claims.

1, wherein said nucleotide sequence encodes a CRFR2, wherein the mouse exhibits a phenotype of decreased activity, hypoactivity, and a decreased susceptibility seizure, and a method of making the same transgenic mouse comprising introducing a targeting construct into an ES cell, introducing the ES cell into a blastocyst, and implanting the blastocyst into a pseudopregnant mouse, and allowing said blastocyst to develop to term, does not reasonably provide enablement for all other transgenic non-human animals and methods of making transgenic mice embraced by the claims. The specification does not enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the invention commensurate in

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The claims are directed transgenic non-human animals, particularly a mouse, that comprise a disruption in a CRFR2 gene, the sequence of which is set forth in SEQ ID NO: 1, wherein the mouse exhibits a phenotype of decreased activity, hypoactivity, and decreased susceptibility to seizures. The claims are further directed a method of producing a transgenic mouse comprising a disruption in a CRFR2 gene.

The specification teaches the generation of transgenic mice by disruption of the nucleotide sequence set forth in SEQ ID NO: 1, wherein SEQ ID NO: 1 encodes a CRFR2. See page 6, at lines 11-17, page 9 at lines 15-24, and the working example on pages 53-54, of the specification. The specification teaches that transgenic mice whose genome comprises a homozygous disruption in SEQ ID NO:1 exhibit a phenotype of decreased activity, hypoactivity, and decreased susceptibility to seizures in response to metrazol as compared to wild-type mice, as a result of the disruption of the nucleotide

sequence set forth in SEQ ID NO: 1. See pages 53-54 of the specification. While the specification has taught the generation of such a transgenic knockout mouse having a phenotype of decreased activity, hypoactivity, and decreased susceptibility to seizures, the specification has not taught the generation of the other transgenic non-human animals encompassed by the claims. The specification has also taught a method of producing a transgenic mouse comprising a disruption in a CRFR2 gene as set forth in the nucleotide sequence set forth in SEQ ID NO: 1, wherein the method requires introduction of a targeting construct into an embryonic stem cell. The specification has not taught how to create a transgenic mouse comprising a disruption in a CRFR2 gene wherein a targeting construct is introduced into any other cell. The working examples, guidance and relevant teachings provided by the instant specification are directed to the creation of the above transgenic mouse but do not support the creation of other transgenic non-human animals encompassed by the claims. See pages 53-54.

With regard to claim breadth, the standard under 112, first paragraph entails the determination of what the claims recite and what the claims mean as a whole. In addition, when analyzing the enabled scope of the claims, the teachings of the specification are taken into account because the claims are to be given their broadest reasonable interpretation that is consistent with the specification. As such, in light of the specification, the claimed invention is properly interpreted with regard to the disclosed phenotype of the exemplified transgenic mouse whose genome comprises a homozygous disruption of the nucleotide sequence set forth in SEQ ID NO: 1. Such an interpretation is consistent with the specification despite that the claimed non-human

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animals require only that they comprise a disrupted CRFR2, particularly the nucleotide sequence set forth in SEQ ID NO: 1. This is because, with regard to the enablement requirement, one of skill in the art must be provided with both how to make and use the claimed invention. As such, the enabled scope of the claimed invention, in light of the teachings of the specification, is found to be the generation of a transgenic mouse whose genome comprises a homozygous disruption of the nucleotide sequence set forth in SEQ ID NO: 1 which exhibit a phenotype of decreased activity, hypoactivity, and decreased susceptibility to seizures.

The following aspect of the rejection under 35 U.S.C. 112, first paragraph is directed to claims 8, 10 and 17-22 as they read on transgenic knockout non-human animals, use of embryonic stem cells to make a transgenic mouse, and germline transmission of ES cells:

Both the specification and the state of the art have taught that the transgenic knockout technology requires the use of embryonic stem cells that have been genetically manipulated to comprise a disruption in a nucleotide sequence of interest. The specification has not taught creation of a transgenic knockout non-human animal by methods that do not require embryonic stem cells. Presently, the transgenic knockout technology is limited to the mouse system. See below.

With regard to the claim breadth directed to transgenic non-human animals, the specification fails to teach the production of any transgenic non-human animal comprising a disruption in a CRFR2 gene other than a transgenic knockout mouse whose genome comprises a homozygous disruption in the nucleotide sequence set

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forth in SEQ ID NO: 1. It is well known in the knockout art that the production of knockout animals other than mice is undeveloped. This is because ES cell technology is generally limited to the mouse system, at present, and that only "putative" ES cells exist for other species. See Moreadith et al. at page 214, Summary. Seamark (Reproductive Fertility and Development, 1994) supports this observation by reporting that totipotency for ES cell technology in many livestock species has not been demonstrated (page 6, Abstract). Likewise, Mullins et al (Journal of Clinical Investigation, 1996) state that "although to date chimeric animals have been generated from several species including the pig, in no species other than the mouse has germline transmission of an ES cell been successfully demonstrated." (page S38, column 1, first paragraph). Moreover, with regard to claim 10 neither the state of the art nor the prior art of record has provided guidance for use of cells, other than ES cells for production of a transgenic knockout mouse. It would be unpredictable if other cells could be used for the production of a transgenic knockout mouse because other cells may be not totipotent or transmit through the germline as ES cells do. Even more, claims 8 and 17-22 as written do not appear to require germline transmission of the disrupted nucleotide sequence. These claims may be broadly interpreted to read on a single cell comprising a disrupted nucleotide sequence. Since the claims do not require germline transmission of the disrupted nucleotide sequence it would be unpredictable if an ES cell comprises the disrupted nucleotide sequence. As stated above the evidence of record does not support germline transmission of non-ES cells. Also, it would be unpredictable if a disruption of a nucleotide sequence in a single cell would result in a phenotype; the

instant specification has not provided any uses for a transgenic mouse that does not exhibit a phenotype resulting from disruption of a nucleotide sequence (see below). As the claims are directed to transgenic non-human animals (claim 8) or a method that requires the use of a cell to in the production of a transgenic mouse (claim 10), wherein the cell is interpreted to read on an embryonic stem cell (as in claim 10) comprising a disruption in a CRFR2 gene, which must be generated by the introduction of a transgene into an ES cell or transgenic non-human animals, particularly a mouse, that do not exhibit germline transmission of a disrupted nucleotide sequence, the state of the art supports that only mouse ES cells were available for use for production of transgenic mice whose genomes comprise a homozygous disruption of a CRFR2 gene as set forth in SEQ ID NO: 1. Given the unpredictable state of the art it would have required undue experimentation for the skilled artisan to create transgenic knockout non-human animals of species other than the mouse or to make a transgenic knockout mouse with a cell other than an embryonic stem cell.

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Claims 8 and 17 encompass transgenic non-human animals, particularly a mouse, that comprise a disruption in a CRFR2 gene, particularly the nucleotide sequence set forth in SEQ ID NO: 1, that do not exhibit any particular phenotype. The state of the art at the time of filing was such that one of skill could not predict the phenotype of a knockout mouse (Moreadith et al., 1997, J. Mol. Med., Vol. 75, pages 208-216; see page 208, column 2, last full paragraph). Moens et al. (Development, Vol. 119, pages 485-499, 1993) disclose that two mutations produced by homologous recombination in two different locations of the N-myc gene produce two different

phenotypes in mouse embryonic stem cells, one leaky and one null (see abstract). The specification has asserted that the nucleotide sequence set forth in SEQ ID NO: 1 encodes a CRFR2. However, it would be difficult to predict any phenotype resulting from disruption of the sequence of SEQ ID NO: 1 in light of the above. The specification discloses a phenotype exhibited by knockout mice comprising a disruption in the nucleotide sequence set forth in SEQ ID NO: 1 is decreased activity, hypoactivity, and decreased susceptibility to seizures. See pages 53-54 of the specification. Claims 8 and 17, as written, do not include a phenotype that differs from the wild-type mouse. Moreover the skilled artisan would not know how to use a transgenic knockout nonhuman animal that lacks a phenotype, particularly because the instant specification has not provided uses for such; the transgenic mice that have a phenotype may be used for drug testing according to the instant specification. The specification overcomes the unpredictability in obtaining a phenotype associated with a disruption of the nucleotide sequence set forth in SEQ ID NO: 1, which is asserted to encode a CRFR2; however, the claims are not commensurate in scope with the enabled phenotype disclosed in the specification. Inclusion of a phenotype associated with a disruption of the nucleotide sequence set forth in SEQ ID NO: 1 or a CRFR2 gene in a mouse in the claims would overcome this aspect of the rejection. Given the unpredictable nature of a phenotype that results from disruption of a nucleotide sequence it would have required undue experimentation for the skilled artisan to use a transgenic non-human knockout animal that lacks a phenotype.

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As a final issue, claims 17-22 encompass transgenic mice comprising a disruption in a homolog of the nucleotide sequence set forth in SEQ ID NO: 1. The specification has disclosed mice that comprise a disruption in the nucleotide sequence set forth in SEQ ID NO: 1, while the specification has not taught mice comprising a disruption in a homolog of the nucleotide sequence set forth in SEQ ID NO: 1. The claims broadly encompass disruption of nucleotide sequences that are homologs of the nucleotide sequence set forth in SEQ ID NO: 1, which have different structures from the nucleotide sequence set forth in SEQ ID NO: 1; given the structural differences it may presumed that the encoded proteins possess different functions. Moreover, since the claims broadly encompass disrupting homologs of SEQ ID NO: 1, the members of the genus of such homologs may possess different functions and chemical structures, it would be unpredictable if disrupting homologs of SEQ ID NO: 1, would result in the phenotype of decreased activity, hypoactivity, and decreased susceptibility to seizures as exhibited by transgenic mouse exemplified in the working example on pages 53-54 of the specification; the specification has not disclosed any homologs of the nucleotide sequence set forth in SEQ ID NO: 1. The issue of the unpredictability of a phenotype resulting from disruption of a homolog of SEQ ID NO: 1 arises because the state of the art at the time of filing was such that one of skill could not predict the phenotype of a knockout mouse as discussed above. See Moreadith. Moens et al. (see above) disclose that two mutations produced by homologous recombination in two different locations of the N-myc gene produce two different phenotypes in mouse embryonic stem cells, one leaky and one null (see abstract). The specification has asserted that

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the nucleotide sequence set forth in SEQ ID NO: 1 encodes a CRFR2 but has not provided any teachings with regard to homologs of the nucleotide sequence set forth in SEQ ID NO: 1. It would be difficult to predict any phenotype resulting from disruption of a homolog of the nucleotide sequence of SEQ ID NO: 1 in light of the above. The specification discloses a phenotype exhibited by knockout mice comprising a disruption in the nucleotide sequence set forth in SEQ ID NO: 1 is decreased activity, hypoactivity, and decreased susceptibility to seizures but has not disclosed a phenotype resulting from the disruption of a homolog of SEQ ID NO: 1. As such it would have required undue experimentation for the skilled artisan to make and use a transgenic mouse comprising a disruption of a homolog of the nucleotide sequence set forth in SEQ ID NO: 1 without a reasonable expectation of success.

Therefore, in view of the quantity of experimentation necessary to determine the parameters listed above for the production of transgenic non-human animals comprising a disruption in a CRFR2 gene, the lack of direction or guidance provided by the specification for the production of transgenic non-human animals comprising a disruption in a CRFR2 gene, the absence of working examples for the demonstration or correlation to the production of a transgenic knockout non-human animal that exhibits a phenotype other than the exemplified mouse, the unpredictable state of the art with respect to a phenotype that results from disruption of a given nucleotide sequence, the undeveloped art pertaining to the establishment of true embryonic stem (ES) cells of animal species other than mouse, and the breadth of the claims drawn to all non-human animals and to homologs of the nucleotide sequence set forth in SEQ ID NO: 1, it would

have required undue experimentation for one skilled in the art to make and/or use the claimed invention.

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It is noted that the following claim language may be sufficient to overcome the preceding enablement rejection: A transgenic mouse whose genome comprises a homozygous disruption of a CRFR2 gene, the nucleotide sequence of which is set forth in SEQ ID NO: 1, exhibiting a phenotype of decreased activity, hypoactivity, and decreased susceptibility to seizures.

Claims 17-22 are rejected under 35 U.S.C. 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to reasonably convey to one skilled in the relevant art that the inventor(s), at the time the application was filed, had possession of the claimed invention.

The claims are directed transgenic non-human animals, particularly a mouse, that comprise a disruption in the nucleotide sequence set forth in SEQ ID NO: 1 or a homolog thereof, wherein the mouse exhibits a phenotype of decreased activity, hypoactivity, and decreased susceptibility to seizures.

Vas-Cath Inc. v. Mahurkar, 19USPQ2d 1111 (Fed. Cir. 1991), clearly states that "applicant must convey with reasonable clarity to those skilled in the art that, as of the filing date sought, he or she was in possession of the invention. The invention is, for purposes of the 'written description' inquiry, whatever is now claimed." Vas-Cath Inc. v. Mahurkar, 19USPQ2d at 1117. The specification does not "clearly allow persons of

ordinary skill in the art to recognize that [he or she] invented what is claimed." <u>Vas-Cath</u> Inc. v. Mahurkar, 19USPQ2d at 1116.

The specification has provided a description for the nucleotide sequence set forth in SEQ ID NO: 1. The specification has asserted that the nucleotide sequence set forth in SEQ ID NO: 1 encodes a CRFR2. However, the nucleotide sequences that homologs of the nucleotide sequence set forth in SEQ ID NO: 1 have not been disclosed. Based upon the prior art there is expected to be variation among the species of polynucleotides that comprise the genus of nucleotide sequences as set forth in SEQ ID NO: 1. The specification has failed to disclose the nucleotide sequences of any nucleic acid that are homologs of the nucleotide sequence set forth in SEQ ID NO: 1. There is no evidence on the record of a relationship between the structures of any DNA molecules, which are homlogs of the nucleotide sequence set forth in SEQ ID NO: 1, that would provide any reliable information about the structures of other such DNA molecules. There is no evidence on the record that the nucleotide sequences that are homologs of the nucleotide sequence set forth in SEQ ID NO: 1 had a known structural relationship to other DNA sequences encompassed within the genus. Furthermore, the evidence of record has not provided evidence of a structural relationship between the nucleotide sequence set forth in SEQ ID NO: 1 and the nucleotide sequences that homologs of the nucleotide sequence set forth in SEQ ID NO: 1. Moreover it is not known if the homologs of SEQ ID NO: 1 would encode proteins that would even possess the biological activity of the protein encoded by the nucleotide sequence set forth in SEQ ID NO: 1. The claimed invention as a whole is not adequately described if



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the claims require essential or critical elements which are not adequately described in the specification and which is not conventional in the art as of applicants effective filing date. Possession may be shown by actual reduction to practice, clear depiction of the invention in a detailed drawing, or by describing the invention with sufficient relevant identifying characteristics such that a person skilled in the art would recognize that the inventor had possession of the claimed invention. <u>Pfaff v. Wells Electronics, Inc.</u>, 48 USPQ2d 1641, 1646 (1998).

In the instant case the claimed embodiments of nucleotide sequence that are homologs of the nucleotide sequence set forth in SEQ ID NO: 1 lack a written description. The specification fails to describe what DNA molecules fall into this genus. The skilled artisan cannot envision the detailed chemical structure of the encompassed DNA molecules that are homologs of the nucleotide sequence forth in SEQ ID NO: 1, and therefore conception is not achieved until reduction to practice has occurred, regardless of the complexity or simplicity of the method of isolation. Adequate written description requires more than a mere statement that it is part of the invention and reference to a potential method of isolating it. See *Fiers v. Revel*, 25 USPQ2d 1601, 1606 (Fed. Cir. 1993) and *Amgen Inc. v. Chugai Pharmaceutical Co. Ltd.*, 18 USPQ2d 1016 (Fed. Cir. 1991).

One cannot describe what one has not conceived. See *Fiddes v. Baird*, 30 USPQ2d 1481, 1483. In *Fiddes*, claims directed to mammalian FGF's were found to be unpatentable due to lack of written description for that broad class. The specification provided only the bovine sequence.

In view of the above considerations one of skill in the art would not recognize that applicant was in possession of the necessary common features or attributes possessed by member of the genus of nucleotide molecules that are homologs of the nucleotide sequence forth in SEQ ID NO: 1. Moreover, the art would generally recognize that there would be variation among the species of the genus of polynucleotide molecules that are homologs of the nucleotide sequence set forth in SEQ ID NO: 1. Therefore, Applicant was not in possession of the genus of nucleotide molecules that are homologs of the nucleotide sequence set forth in SEQ ID NO: 1 as encompassed by the claims. University of California v. Eli Lilly and Co., 43 USPQ2d 1398, 1404, 1405 held that to fulfill the written description requirement, a patent specification must describe an invention and do so in sufficient detail that one skilled in the art can clearly conclude that "the inventor invented the claimed invention."

## Claim Rejections - 35 USC § 102

The following is a quotation of the appropriate paragraphs of 35 U.S.C. 102 that form the basis for the rejections under this section made in this Office action:

A person shall be entitled to a patent unless -

- (a) the invention was known or used by others in this country, or patented or described in a printed publication in this or a foreign country, before the invention thereof by the applicant for a patent.
- (e) the invention was described in-

(2) a patent granted on an application for patent by another filed in the United States before the invention by the applicant for patent, except that a patent shall not be deemed filed in the United States for the purposes of this subsection based on the filing of an international application filed under the treaty defined in section 351(a).

<sup>(1)</sup> an application for patent, published under section 122(b), by another filed in the United States before the invention by the applicant for patent, except that an international application filed under the treaty defined in section 351(a) shall have the effect under this subsection of a national application published under section 122(b) only if the international application designating the United States was published under Article 21(2)(a) of such treaty in the English language; or

Claims 8 and 10 are rejected under 35 U.S.C. 102(a) as being anticipated by Coste et al (Nature Genetics, 2000, 24: 403-409).

The claims are directed to a transgenic non-human animal comprising a disruption in a CRFR2 gene and a method for producing a transgenic mouse comprising a disruption in a CRFR2 gene.

For the purposes of the this rejection a CRFR2 gene is interpreted to be a CRHR2. This interpretation has been made because the prior art as set forth in the specification on pages 1-4 sets forth that a CRFR2 gene and a CRHR2 gene are the same. The difference being in name only, corticotropin-releasing factor receptor as opposed to corticotropin-releasing hormone receptor.

Coste et al teach a transgenic mouse comprising a disruption in the CRHR2 gene. Coste et al teach that the transgenic mouse is created by introducing a targeting vector into ES cells, transferring the ES cells to a blastocyst and then implanting the blastocyst into a pseudopregnant female mouse, wherein said female mouse gives birth to a chimeric mouse, and breeding said chimeric mouse to produce the transgenic mouse. See page 403, column 1, first paragraph as well as the Materials and Methods section on pages 406-408.

Thus, the teachings of Coste et al anticipate all of the instant claim limitations.

Claims 8 and 10 are rejected under 35 U.S.C. 102(a) as being anticipated by Bale et al (Nature Genetics, 2000, 24: 410-414).

The claims are directed to a transgenic non-human animal comprising a disruption in a CRFR2 gene and a method for producing a transgenic mouse comprising a disruption in a CRFR2 gene.

For the purposes of the this rejection a CRFR2 gene is interpreted to be a CRHR2. This interpretation has been made because the prior art as set forth in the specification on pages 1-4 sets forth that a CRFR2 gene and a CRHR2 gene are the same. The difference being in name only, corticotropin-releasing factor receptor as opposed to corticotropin-releasing hormone receptor.

Bale et al teach a transgenic mouse comprising a disruption in the CRHR2 gene. Coste et al teach that the transgenic mouse is created by introducing a targeting vector into ES cells, transferring the ES cells to a blastocyst and then implanting the blastocyst into a pseudopregnant female mouse, wherein said female mouse gives birth to a chimeric mouse, and breeding said chimeric mouse to produce the transgenic mouse. See page 410, column 1, first paragraph bridging to page 411 as well as the Materials and Methods section on page 412.

Thus, the teachings of Bale et al anticipate all of the instant claim limitations.

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Claims 8 and 10 are rejected under 35 U.S.C. 102(a) as being anticipated by Kishimoto et al (Nature Genetics, 2000, 24: 415-419).

The claims are directed to a transgenic non-human animal comprising a disruption in a CRFR2 gene and a method for producing a transgenic mouse comprising a disruption in a CRFR2 gene.

For the purposes of the this rejection a CRFR2 gene is interpreted to be a CRHR2. This interpretation has been made because the prior art as set forth in the specification on pages 1-4 sets forth that a CRFR2 gene and a CRHR2 gene are the same. The difference being in name only, corticotropin-releasing factor receptor as opposed to corticotropin-releasing hormone receptor.

Kishimoto et al teach a transgenic mouse comprising a disruption in the CRHR2 gene. Kishimoto et al teach that the transgenic mouse is created by introducing a targeting vector into ES cells, transferring the ES cells to a blastocyst and then implanting the blastocyst into a pseudopregnant female mouse, wherein said female mouse gives birth to a chimeric mouse, and breeding said chimeric mouse to produce the transgenic mouse. See page 415, column 1, first paragraph, Figure 1 on page 415 as well as the Materials and Methods section on pages 418.

Thus, the teachings of Kishimoto et al anticipate all of the instant claim limitations.

Claims 8 and 10 are rejected under 35 U.S.C. 102(e) as being anticipated by Lee et al (US 6,353,152; effective filing date of 7/15/1999).

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The claims are directed to a transgenic non-human animal comprising a disruption in a CRFR2 gene and a method for producing a transgenic mouse comprising a disruption in a CRFR2 gene.

Lee et al teach a transgenic mouse comprising a disruption in the CRHF2 gene.

Lee et al teach that the transgenic mouse is created by introducing a targeting vector into ES cells, transferring the ES cells to a blastocyst and then implanting the blastocyst into a pseudopregnant female mouse, wherein said female mouse gives birth to a chimeric mouse, and breeding said chimeric mouse to produce the transgenic mouse. See column 8 and throughout entire document.

Thus, the teachings of Lee et al anticipate all of the instant claim limitations.

#### Conclusion

No claim is allowed. Claims 17-22 appear to be free of the prior art of record but are subject to other rejections.

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Any inquiry concerning this communication or earlier communications from the examiner(s) should be directed to Peter Paras, Jr., whose telephone number is 703-308-8340. The examiner can normally be reached Monday-Friday from 8:30 to 4:30 (Eastern time).

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Deborah Reynolds, can be reached at 703-305-4051. Papers related to this application may be submitted by facsimile transmission. Papers should be faxed via the PTO Fax Center located in Crystal Mall 1. The faxing of such papers must conform with the notice published in the Official Gazette, 1096 OG 30 (November 15, 1989). The CM1 Fax Center numbers are (703) 308-4242 and (703) 305-3014.

Inquiries of a general nature or relating to the status of the application should be directed to Patsy Zimmerman whose telephone number is (703) 308-0009.

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Peter Paras, Jr.

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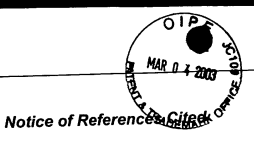
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				U.S. PATENT DOCUMENTS	
*		Document Number Country Code-Number-Kind Code	Date MM-YYYY	Name	Classification 800/18
	Α	US-6353152 B1	03-2002	Lee et al	800/16
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	U	Moens et al. Defects in heart and lung development in compound heterozygotes for two different targeted mutations at the N-myc locus. Development, 1993, 119: 485-499.
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TECH CENT S Page 1 and Total amark Office; U.S. DEPARTMENT OF COMMERCE the Paperwork Reduction Act of 1995, no persons are required to essential total control number. Complete if Known FEE TRANSMITTAL 09/900,700 Application Number Filing Date July 6, 2001 for FY 2003 Keith D. Allen First Named Inventor Effective 01/01/2003. Patent fees are subject to annual revision. **Examiner Name** Peter Paras Jr. Applicant claims small entity status. See 37 CFR 1.27 1632 Art Unit (\$) 55.00 **TOTAL AMOUNT OF PAYMENT** R-616 Attorney Docket No. METHOD OF PAYMENT (check all that apply) FEE CALCULATION (continued) Money 3. ADDITIONAL FEES Check Credit card Other None Large Entity L Small Entity Deposit Account: Fee Fee **Fee Description** Deposit Code Code (\$) (\$) Fee Paid 50-1271 Account 2051 1051 130 65 Surcharge - late filing fee or oath Number Deposit 2052 Surcharge - late provisional filing fee or 1052 50 Deltagen, Inc. Account cover sheet Name 1053 130 1053 130 Non-English specification The Commissioner is authorized to: (check all that apply) 1812 2,520 For filing a request for ex parte reexamination 1812 2,520 ✓ Charge fee(s) indicated below Credit any overpayments 920\* Requesting publication of SIR prior to 1804 920 1804 Charge any additional fee(s) during the pendency of this application Examiner action Charge fee(s) indicated below, except for the filing fee Requesting publication of SIR after 1805 1,840 1805 1.840° Examiner action to the above-identified deposit account. 1251 110 2251 Extension for reply within first month 55 55.00 **FEE CALCULATION** Extension for reply within second month 1252 410 2252 205 1. BASIC FILING FEE 1253 930 2253 Extension for reply within third month arge Entity Small Entity Fee Description Fee Paid Fee ree Code (\$) 2254 1254 1.450 725 Extension for reply within fourth month Code (\$) 985 Extension for reply within fifth month 2255 1255 1.970 1001 750 2001 375 Utility filing fee 2401 1401 320 1002 330 2002 160 Notice of Appeal 165 Design filing fee 2402 160 Filing a brief in support of an appeal 1402 320 1003 520 2003 260 Plant filing fee 140 Request for oral hearing 2403 1004 750 2004 375 Reissue filing fee 1403 280 1005 160 2005 80 Provisional filing fee 1451 1,510 1451 1,510 Petition to institute a public use proceeding 2452 55 Petition to revive - unavoidable 110 1452 SUBTOTAL (1) (\$) 1453 1,300 2453 650 Petition to revive - unintentional 2. EXTRA CLAIM FEES FOR UTILITY AND REISSUE 1501 1,300 2501 650 Utility issue fee (or reissue) Fee from Extra Claims Fee Paid below 1502 470 2502 235 Design issue fee **Total Claims** X 1503 630 2503 315 Plant issue fee Independent 1460 1460 130 130 Petitions to the Commissioner Multiple Dependent 1807 50 1807 50 Processing fee under 37 CFR 1.17(q) arge Entity Small Entity 180 180 Submission of Information Disclosure Stmt 1806 1806 Fee Description 40 Recording each patent assignment per Code (\$) Code (\$) 8021 40 8021 property (times number of properties) Claims in excess of 20 1202 18 2202 9 375 Filing a submission after final rejection (37 CFR 1.129(a)) 1809 750 2809 Independent claims in excess of 3 1201 84 2201 42 1203 280 2203 140 Multiple dependent claim, if not paid 1810 750 2810 375 For each additional invention to be examined (37 CFR 1.129(b)) \*\* Reissue independent claims 1204 2204 over original patent 1801 750 2801 375 Request for Continued Examination (RCE) 1802 900 1802 900 Request for expedited examination 1205 18 2205 9 \*\* Reissue claims in excess of 20 and over original patent of a design application Other fee (specify) (\$) SUBTOTAL (2) \*Reduced by Basic Filing Fee Paid

SUBTOTAL (3) (\$) 55.00 \*\*or number previously paid, if greater; For Reissues, see above SUBMITTED BY (Complete (if applicable) Registration No. Aaron T. Hokamura Telephone 650-569-5171 Name (Print/Type) 51.810 (Attorney/Agent) the Signature WW Date 02/24/03

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This collection of information is required by 37 CFR 1.17 and 1.27. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 12 minutes to complete, including gathering, preparing, and submitting the completed application form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, Washington, DC 20231. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Commissioner for Patents, Washington, DC 20231.